# IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of:

Confirmation No: 2134

FOSTER et al.

Art Unit: 1654

Appl. No.: 09/937,484

Examiner: Audet, Maury A.

§ 371 Date: January 23, 2002

Atty. Docket: 1581.0870000/TJS/LDB

For: Use of a Lectin or Conjugates for Modulation of C-Fibre Activity

Declaration Under 37 C.F.R. § 1.132 of Dr. John Chaddock

Assistant Commissioner for Patents Washington, D.C. 20231

Sir:

I, the undersigned, Dr. John Chaddock, do hereby solemnly and sincerely declare that:

- 1. I am a named inventor of the subject matter described and claimed in U.S. Patent Application No. 09/937,484.
- 2. I am currently Head of Molecular Biology at Syntaxin, Ltd. in the United Kingdom. As evidenced by the abridged version of my curriculum vitae (attached), I have been actively undertaking research in the field of lectin conjugates for the last 18 years. I am an expert in the field of modulating C-fibre neuron activity for therapeutic purposes.
- 3. I am familiar with the Office Action dated 29 November 2005 for the above-captioned case.

- 4. Lectin compounds are a well-recognized group of structurally and functionally related molecules that share common structural features. As a first structural feature, lectins possess a highly conserved binding site triad of amino acids, which is known as the "Asp-Gly-Asn triad". The conserved Asp-Gly-Asn triad is disclosed, for example, in Svensson et al. J. Mol. Biol. 321: 69-83 (2002) (enclosed herein as Exhibit A), and in Loris et al. J. Mol. Biol. 335: 1227-1240 (2004) (enclosed herein as Exhibit B).
- 5. As a second structural feature, the lectins possess a "lectin fold", which consists primarily of three  $\beta$ -sheets:
  - a "flat" six-membered "back" β-sheet;
  - a small "top" β-sheet; and
  - a curved, seven-stranded "front" β-sheet.

The "lectin fold" is described in detail in Chandra et al. Prot. Engin. 14: 857-866 (2001) (enclosed herein as Exhibit C), and in Turton et al. Glycobiology 14: 923-929 (2004) (enclosed herein as Exhibit D). The "lectin fold" facilitates presentation of a substrate (sugar) to the lectin in a concave region of the "front"  $\beta$ -sheet. The highly conserved amino acid triad consisting of Asp, Asn and Gly is found within the "lectin fold."

- 6. The primary sequence of Erythrina cristagalli lectin (ECL) is disclosed in Figure 2 of Exhibit A. Alignment of the primary sequence of ECL with the sequence of the Erythrina corallodendron lectin (ECorL) in the same Figure 2 of Exhibit A shows that ECL shares 96% sequence homology with ECorL, and reveals 100% identity in the conserved Asp-Gly-Asn triad: Asp89, Gly107 and Asn 133.
- 7. The primary sequence of the first 244 amino acids of the Erythrina corallodendron lectin (ECorL) was first disclosed in Adar et al. FEBS Lett. 257: 81-85 (1989) (enclosed herein as Exhibit E). Comparison of the amino acid sequence of ECorL with that of other legume lectins

reveals a high degree of homology overall and 100% identity in the conserved Asp-Gly-Asn triad. See Figure 3 in Exhibit E. This highly conserved lectin binding site "Asp-Gly-Asn triad" is well known to those skilled in the art.

8. It is understood by researchers in the field of lectins that the Asp-Gly-Asn triad and the lectin fold are highly conserved distinguishing features of all lectins.

In this regard, Loris R. et al., Proteins, 1994 Dec; 20(4):330-346 (enclosed herein as Exhibit F) describes monosaccharide recognition by two forms of lentil lectin, via a highly conserved triad of residues, namely Asp 81, Gly 99 and Asn 125. Thomas, C.J. and Surolia A., Biochem. Biophys. Res. Commun 2000 Feb 16; 268 (enclosed herein as Exhibit G) confirms that an invariant triad of residues Asp 87, Gly 105 and Asn 137 coordinates recognition of L-fucose by fucose-binding legume lectins. These references confirm that the "Asp-Gly-Asn triad" is essential for ligand recognition in a variety of lectins whose crystal structures are known.

Rao VS. et al., J. Biomol. Struct. Dyn 1998 Apr 15(5):853-860 (enclosed herein as Exhibit H) confirms, with reference to soybean agglutinin, that the invariant "Asp-Gly-Asn triad" is essential for binding carbohydrate. Together with an aromatic residue (Phe or Tyr), these three invariant residues provide the basic frame for the sugar to bind. This is confirmed by Rao VS. et al., Int. J. Macromol. 1998 Nov; 23(4):295-307 (enclosed herein as Exhibit I), which reviews the sugar binding sites of Erythrina corallodendron (EcorL), peanut lectin (PNA), Lathyrus ochrus (LOLI) and pea lectin (PSL) and confirms that the invariant residue Asp (from loop A), the invariant residue Asn (from loop C) and the invariant residue Gly (from loop B) are required for a tight interaction with the sugar molecule.

8. I further state that all statements made on my own knowledge are true and that all statements made on information and belief are believed to be true and further that willful false

statements and the like are punishable by fine or imprisonment or both, under Section 1001 of Title 18 of the U.S. Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereupon.

Date: 21 st fort 2006 Alchan John Chaddock, Ph.D.

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#### **BIOGRAPHICAL SKETCH**

Provide the following information for the key personnel in the order listed for Form Page 2. Follow this format for each person, DO NOT EXCEED FOUR PAGES.

Head of Molecular Biology

NAME POSITION TITLE

INSTITUTION AND LOCATION	DEGRÉÉ (if applicable)	YEAR(\$)	FIELD OF STUDY
University of Durham, UK	BSc.	1988	Molecular Biology / Biochemistry
University of Warwick, UK	Ph.D.	1992	Protein Biochemistry
Open University, UK	Cert. Mgmt	2003	Management

NOTE: The Biographical Sketch may not exceed four pages, Items A and B (together) may not exceed two of the four-page limit. Follow the formats and instructions on the attached sample.

A. Positions and Honors. List in chronological order previous positions, concluding with your present position. List any honors. Include present membership on any Federal Government public advisory committee.

#### Positions and Employment

Chaddock, John Andrew

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1992-1995	Postdoctoral Research, University of Warwick, UK
1995-1996	Postdoctoral Research, University of Warwick, UK
1996-2001	Scientist, Centre for Applied Microbiology & Research, Salisbury, UK
2002-2005	Senior Scientist, Health Protection Agency, Centre for Applied Microbiology & Research, Salisbury, UK
2005-2006	Head of Molecular Biology, Syntaxin Limited, Salisbury, UK

## Other Experience

MSc Examiner in the field of protein toxin biochemistry Co-supervisor of Ph.D. studentship at the University of Bath Visiting lecturer at University of Bath

#### **Professional Memberships**

1988-present Member of Biochemical Society, UK
2002-2004 Member of International Association for the Study of Pain

## B. Selected peer-reviewed publications (in chronological order).

- Wales, R. Chaddock, J. A., Roberts, L. M. & Lord, J. M. (1992) Addition of an ER retention signal to the ricin Achain increases the cytotoxicity of the holotoxin. Exp. Cell Res., 203, 1-4.
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- Chaddock, J. A. and Melling, J. (2002) Clostridium botulinum and associated neurotoxins. Chapter 55. Molecular Medical Microbiology. Ed Max Sussman, Academic Press, London. 1141-1152.
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dol:10.1016/50022-2636(02)00554-5 available online at http://www.kleslibrary.com on IBE

J. Mol. Biol. (2002) \$21, 69-83



# High-resolution Crystal Structures of Erythrina cristagalli Lectin in Complex with Lactose and 2'-α-L-Fucosyllactose and Correlation with Thermodynamic Binding Data

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The primary sequence of Erythrina cristagalli lectin (ECL) was mapped by mass spectrometry, and the crystal structures of the lectin in complex with lactose and 2-a-1-fucosyllactose were determined at 1.6 Å and 1.7 Å resolution, respectively. The two complexes were compared with the crystal structure of the closely related Erythrina corallodendron lectin (BCorL) in complex with lactose, with the crystal structure of the Ulex europaeus lectin II in complex with 2'-a-L-fucosyllactose, and with two modeled complexes of BCorL with 2'-a-L-fucosyl-N-acetyllactosamine. The molecular models are very similar to the crystal structure of ECL in complex with 2-\alpha-1-fucosyllactose with respect to the overall mode of binding, with the 1-fucose fitting snugly into the cavity surrounded by Tyr106, Tyr108, Trp135 and Pro134 adjoining the primary combining site of the lectin. Marked differences were however noted between the models and the experimental structure in the network of hydrogen bonds and by through the respective holding the 1-fucosyline the respective forms. hydrophobic interactions holding the L-fucose in the combining site of the lectin, pointing to limitations of the modeling approach. In addition to the structural characterization of the ECL complexes, an effort was undertaken to correlate the structural data with thermodynamic data obtained from microcalorimetry, revealing the importance of the water network in the lectin combining site for carbohydrate binding.

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Keywords: crystal structure; glycobiology; lectin; protein-carbohydrate interactions; structure/function

### Introduction

Specific recognition of carbohydrates by proteins lies at the heart of many biological processes, ranging from cell-cell interaction and adhesion of infectious agents to host cells, cellular signaling

Abbreviations used: ECL, Erythrina cristagalli lectin; ECorl, Erythrine corollondendron lectin; ELLA, enzyme-ECarl, Erythrina corallondendron lectin; ELLA, enzyme-linked lectin assay; Fuc, L-fucose; fucosyllactose, Fuclac, 2-a-t-fucosyl-N-acetyllactosamine; Cal, galactose; Glc, glucose; ITC, laothermal titration calorimetry; Lac, lactose; MFD, 2-methyl-2,4-pentanediol; MS, mass spectrometry; rms, root mean square; PBS, phosphate-buffered saline; UEA-II, Lilex europeaus lectin II, WBA-I and II, winged bean acidic lectin I and II. E-mail address of the corresponding author: ute krencel@mobblotech.chalmers.se

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and differentiation, malignancy and metastasis, to fertilization and immune response.12 Understanding the roles of carbohydrates in these processes and how they interact with proteins is expected to have a large impact on the development of new

treatments against many human diseases Legume lectins are especially well suited as a model system to study the molecular basis of protein-carbohydrate recognition because they are structurally similar and yet their specificity is diverse. 1-3 By now, the three dimensional structures of 20 legume lectins have been solved by high-resolution X-ray crystallography. both in few for and in ution X-ray crystallography, both in free form and in complex with a variety of carbohydrate ligandst.

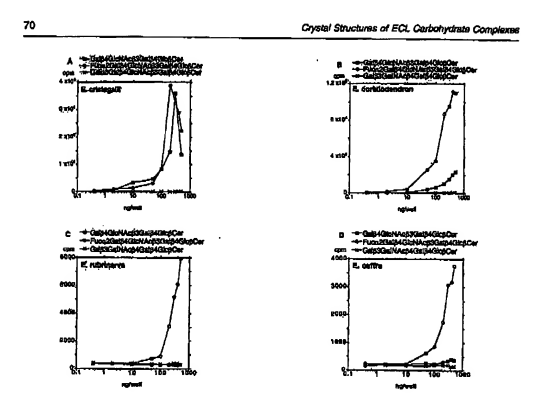
†See 3D Lectin Data Bank on World Wide Web URL: http://www.cermav.cnrs.fr/databank/lectine

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